

Taxon sampling and the accuracy of phylogenetic analyses

Tracy A. HEATH* Shannon M. HEDTKE* David M. HILLIS*,**

(Section of Integrative Biology and Center for Computational Biology and Bioinformatics, One University Station C0930,
The University of Texas, Austin, TX 78712, USA)

Abstract Appropriate and extensive taxon sampling is one of the most important determinants of accurate phylogenetic estimation. In addition, accuracy of inferences about evolutionary processes obtained from phylogenetic analyses is improved significantly by thorough taxon sampling efforts. Many recent efforts to improve phylogenetic estimates have focused instead on increasing sequence length or the number of overall characters in the analysis, and this often does have a beneficial effect on the accuracy of phylogenetic analyses. However, phylogenetic analyses of few taxa (but each represented by many characters) can be subject to strong systematic biases, which in turn produce high measures of repeatability (such as bootstrap proportions) in support of incorrect or misleading phylogenetic results. Thus, it is important for phylogeneticists to consider both the sampling of taxa, as well as the sampling of characters, in designing phylogenetic studies. Taxon sampling also improves estimates of evolutionary parameters derived from phylogenetic trees, and is thus important for improved applications of phylogenetic analyses. Analysis of sensitivity to taxon inclusion, the possible effects of long-branch attraction, and sensitivity of parameter estimation for model-based methods should be a part of any careful and thorough phylogenetic analysis. Furthermore, recent improvements in phylogenetic algorithms and in computational power have removed many constraints on analyzing large, thoroughly sampled data sets. Thorough taxon sampling is thus one of the most practical ways to improve the accuracy of phylogenetic estimates, as well as the accuracy of biological inferences that are based on these phylogenetic trees.

Key words consistency, long-branch attraction, phylogenetic accuracy, phylogenomics, systematic error, taxon sampling, Tree of Life.

The past two decades have seen great progress in reconstructing the Tree of Life. The endeavor of inferring the relationships among all living things is not only of intrinsic interest to biologists, but also has many practical applications throughout biology. Phylogenetic trees allow biologists to make predictions about biology, because we can infer when and where various structures, molecules, or behaviors have evolved in living organisms. Trees also provide information about the expected distribution of these features across taxonomic groups. Moreover, phylogenies facilitate the interpretation of comparative observations by accounting for the historical non-independence of organisms when analyzing across various levels of biological organization (e.g., genes, genomes, individuals, populations, species, or clades).

The number of practical applications of phylogenetics continues to grow each year. For example, phylogenetics has become crucial for comparing and interpreting the various genome sequencing projects.

The very reason that many such projects are undertaken is to provide a broad evolutionary spectrum for interpretation of genome function and evolution in the framework of the Tree of Life (Eisen, 1998). Without a phylogenetic framework, every genome would be a new independent mystery, and the detection of gene function would be greatly hindered without comparative analyses. In the absence of phylogenetic comparisons, studying many gene functions in the human genome would require experimentation and manipulation that is not practical or ethical in humans. Therefore, in a very real sense, the Tree of Life helps us to understand how humans function and how we differ from one another and from other species. The same is also true of all the organisms that we eat, of all the organisms that make us sick, of all the organisms that maintain our ecosystems, and of all the organisms that make the biological world interesting, entertaining, and beautiful.

Applications of phylogenetics are by no means limited to the functional and structural study of genomes, however. Phylogenetic applications span much of biology, from human health (Bush et al., 1999) and forensics (Hillis & Huelsenbeck, 1994; Metzker et al., 2002) to conservation biology (Crandall et al., 2000)

Received: 2 February 2008 Accepted: 23 April 2008

* All authors contributed equally to this work.

** E-mail: dhillis@mail.utexas.edu; phone: 512-471-5792.

and studies of behavior (Martins, 1996). Many of these applications require accurate phylogenetic estimates, not only in terms of tree topologies, but also in branch lengths (for estimation of time and/or the amount of change), ancestral character states (for estimation of evolutionary transitions), and parameters of evolutionary models (for study of evolutionary processes). In general, accurate molecular phylogenetic estimates (estimates that represent true historical relationships among species) are dependent on four primary factors (Swofford et al., 1996): (1) appropriate selection of target genes for analysis; (2) collection of enough sequence data to obtain a robust and repeatable estimate; (3) use of accurate analytical methods; and (4) sufficient taxon sampling for the problem of interest. The first three of these factors often receive the greatest attention from investigators, but increased taxon sampling can be one of the most practical and feasible approaches for improving phylogenetic estimates (Zwickl & Hillis, 2002). Here we explore and review the effects of taxon sampling on phylogenetic analyses and their applications.

1 Dense taxon sampling improves phylogenetic accuracy

Phylogeneticists have long acknowledged that data sets containing a large number of taxa create a more complex computational problem for phylogenetic analysis. As more taxa are added to a phylogenetic data set, the number of possible tree topologies increases very rapidly. In addition, the degree of homoplasy (convergent changes or reversals) increases with the number of taxa (Sanderson & Donoghue, 1989). Regardless, numerous studies on the importance of dense taxon sampling have indicated that introducing additional taxa into a phylogenetic analysis results (on average) in more accurate estimates of evolutionary relationships (Lecointre et al., 1993; Philippe & Douzery, 1994; Hillis, 1996, 1998; Graybeal, 1998; Rannala et al., 1998; Zwickl & Hillis, 2002; Pollock et al., 2002; Poe, 1998a, 2003; DeBry, 2005; Hedtke et al., 2006). These studies represent a broad range of approaches including simulations, examinations of well-studied biological groups, and comparisons to known phylogenies. Each of these approaches has distinct advantages and disadvantages (Hillis, 1995) and together they provide a strong and consistent message about the importance of dense taxon sampling. The benefits of denser taxon sampling are especially evident in conjunction with more thorough searches of solution space (Fig. 1).

Additionally, evaluations of phylogenetic analyses often attribute problematic reconstruction and low resolution to inadequate taxon sampling (e.g., Bremer et al., 1999; Johnson, 2001; Lin et al., 2002; Braun & Kimball, 2002; Chen et al., 2003; Freudenstein et al., 2003; Sorenson et al., 2003; Albrecht et al., 2007).

Although the importance of taxonomic sampling has been intensely investigated, many studies have focused primarily on parsimony and distance methods. Felsenstein (1978) demonstrated that under certain circumstances, parsimony methods are inconsistent, meaning they converge on an incorrect topology as more and more characters are added for a limited sample of taxa. When two non-adjacent taxa share many homoplastic character states along long branches, parsimony methods often interpret such similarity as homology. The resulting tree depicts the two taxa as sister to one another, attributing the shared changes to a branch joining them; this effect is termed long-branch attraction (LBA). Inconsistency is not restricted to parsimony, however, as all phylogenetic reconstruction methods can exhibit this behavior if their assumptions are seriously violated or if there are not enough taxa in the analysis to accurately estimate the parameters of the evolutionary model (Felsenstein, 1978; Hendy & Penny, 1989; DeBry, 1992; Huelsenbeck & Hillis, 1993; Yang, 1994; Huelsenbeck, 1995;

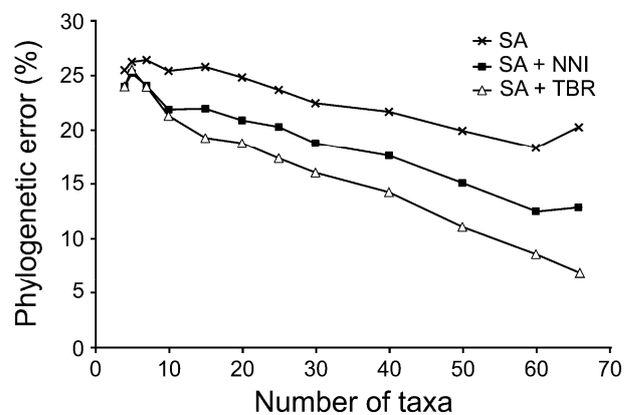


Fig. 1. Error in phylogenetic reconstruction typically decreases with increased taxon sampling of a given taxonomic group. The benefits of increased taxon sampling are particularly evident when searches of the solution space are more thorough. In this graph (adapted from Zwickl & Hillis, 2002, fig. 6), phylogenetic error decreases with increased taxon sampling across all analyses. However, the benefits of adding additional taxa are smaller if only the stepwise-addition algorithm (*SA*) is used to find an approximate solution, compared to the more thorough searches provided by stepwise-addition plus nearest-neighbor-interchanges branch-swapping (*SA+NNI*) or tree bisection-reconnection branch-swapping (*SA+TBR*). Analyses of larger data sets generally require more thorough search algorithms (and thus more computational effort), but result in greatly decreased phylogenetic error.

Lockhart et al., 1996; Gascuel et al., 2001; Huelsenbeck & Lander, 2003; Susko et al., 2004; Philippe et al., 2005). For example, maximum likelihood estimation has been shown to be inconsistent in the presence of severe branch-length heterogeneity (heterotachy, a form of non-stationarity) if the substitution process is assumed to be homogeneous across all lineages (Kolaczkowski & Thornton, 2004; Spencer et al., 2005; Philippe et al., 2005). This example emphasizes the need for probabilistic models that incorporate complex evolutionary processes (Yang & Roberts, 1995; Galtier & Gouy, 1998; Foster, 2004; Blanquart & Lartillot, 2006; Gowri-Shankar & Rattray, 2007; Blanquart & Lartillot, 2008; Kolaczkowski & Thornton, 2008).

Including additional taxa in a phylogenetic analysis will increase the accuracy of the inferred topology by dispersing homoplasy across the tree and reducing the effect of long-branch attraction. Hillis (1996) analyzed data simulated on a 228-taxon tree and showed that simple parsimony and distance methods accurately reconstruct the true topology when provided with sequences 5,000 nucleotides in length. At the time, this result was surprising because it seemingly contradicted the common belief that accurate phylogenetic reconstruction from very large data sets was infeasible. Moreover, Hillis et al. (1994b) had previously shown that analyses of much smaller data sets, containing only 4 taxa, required considerably longer sequences to attain the same level of accuracy. The results of Hillis's (1996) large-scale simulation indicated that for phylogenies containing many taxa,

convergent substitutions or reversals (homoplasy) are distributed among the many lineages in the tree and therefore such misleading information is less likely to overwhelm the true phylogenetic signal.

Because inadequate species sampling can result in trees containing relatively long terminal branches, sparsely sampled data sets are more likely to be affected by LBA. Rannala et al. (1998) simulated ultrametric trees under a simple model of cladogenesis to investigate the impact of removing ingroup taxa on the distribution of branch lengths. They demonstrated that decreasing the proportion of sampled taxa leads to an increase in the average length of terminal branches and generates tree shapes that may be susceptible to long-branch attraction (Fig. 2). Huelsenbeck and Lander (2003) simulated sequences using simple evolutionary models and determined that the probability that parsimony is inconsistent becomes greater as the proportion of taxa sampled decreases and substitution rates increase. Even under very simple models of evolution, unweighted parsimony underestimated the number of changes along branches and converged on an incorrect topology (Huelsenbeck & Lander, 2003).

In general, many studies have shown that adding taxa to bisect long branches can mitigate the effect of LBA (Hendy & Penny, 1989; Graybeal, 1998; Poe & Swofford, 1999; Poe, 2003). However, taxon addition should be practiced judiciously to ensure that enough taxa are added to sufficiently partition multiple long branches (Graybeal, 1998; Poe, 2003) and that the new taxa do not introduce additional long branches (Kim, 1998). Prudent taxon addition is particularly

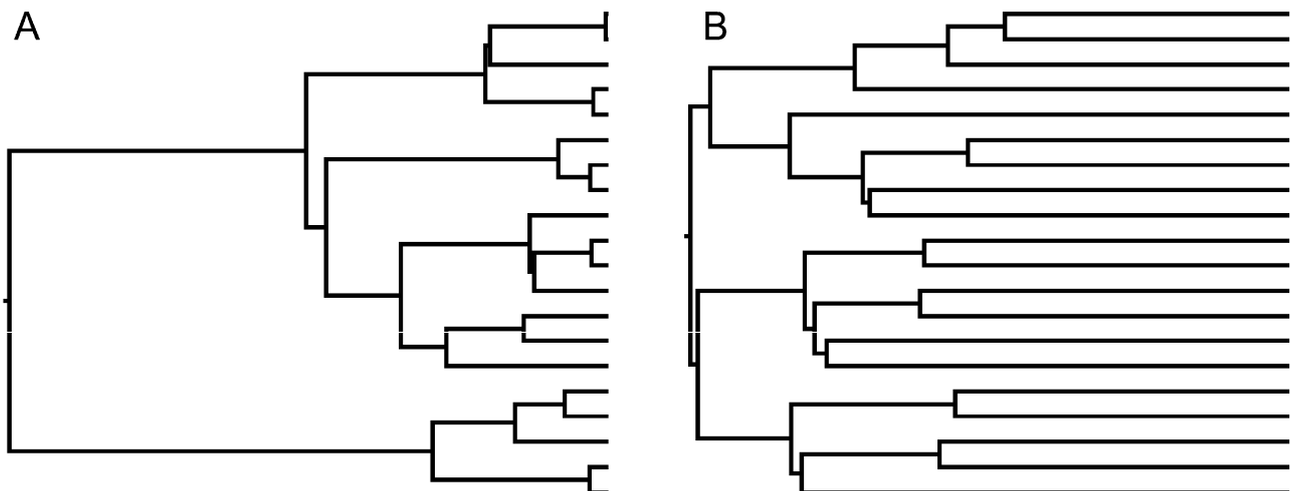


Fig. 2. Two simulations of a birth-death process to model cladogenesis. The speciation rate (λ) and extinction rate (μ) were fixed throughout the simulation and arbitrarily set to $\lambda/\mu=2$. **A**, Phylogenetic tree with complete (100%) taxon sampling (20 taxa total). **B**, Phylogenetic tree with 10% taxon sampling (20 taxa sampled from 200 taxa total). When taxon sampling is low, terminal branch lengths are longer, indicating that sparsely sampled data sets are susceptible to the effects of long-branch attraction. Adapted from Rannala et al. (1998; figs. 1 and 2).

important when conducting parsimony analyses since this method is especially liable to inconsistency due to long-branch attraction. Because parametric methods, such as maximum likelihood, incorporate models that account for unobserved substitutions, these methods are less prone to the effects of long-branch attraction, as long as the models of evolution are adequate. However, enough taxa must be sampled to parameterize these models effectively (Pollock et al., 2002). In addition, longer branches require more accurate models of evolution (because more unobserved changes must be inferred), so increased taxon sampling (which breaks up long branches) greatly benefits parametric methods as well as nonparametric methods. We discuss methods for detecting and minimizing LBA more completely below.

Apart from its effect on topological accuracy, the density of taxon sampling also has an impact on branch-length estimation. Branch lengths provide important information about the amount of change that has occurred over the tree and are critical for applications using phylogenies to make inferences about evolution. Under the parsimony criterion, branch lengths are often underestimated in sparsely sampled regions of the tree because less information is available to infer the history of unobserved substitutions (Fitch & Bruschi, 1987; Fitch & Beintema, 1990). This artifact has been termed the node-density effect (NDE) and may mislead studies that investigate correlations between rates of molecular evolution and biodiversity (Webster et al., 2003; Venditti et al., 2006; Hugall & Lee, 2007). Maximum likelihood, Bayesian, and distance methods are also susceptible to node-density effects, particularly when the assumed model of sequence evolution is overly simple and substitution rates are high (Gojobori et al., 1982; Bruno & Halpern, 1999; Hugall & Lee, 2007). If the density of taxon sampling is increased, additional internal nodes can reveal undetected substitutions and improve estimates of branch lengths.

It has been shown that mis-estimation of branch lengths can, in turn, lead to biased tree topologies (Xia, 2006). Errors in estimates of genetic distance become greater as the amount of divergence between two sequences increases. Pairwise distance methods for phylogenetic reconstruction typically use log-transformed formulae to account for unobserved substitutions (Swofford et al., 1996; Hoyle & Higgs, 2003). When using log-transformed formulae to calculate genetic distances, particularly at high levels of sequence divergence, there is a significant probability that the distance estimates will be undefined

even if the “true” model of sequence evolution is assumed (Hoyle & Higgs, 2003). Therefore, when conducting distance-based analyses, it is very important to consider how taxa are sampled and avoid inclusion of highly divergent sequences.

1.1 Increased taxon sampling versus increased sequence length

Increasing the total number of characters in a data set can increase resolution and support for a phylogeny (Hillis et al., 1994a; Graybeal, 1994; Rannala et al., 1998). In particular, increasing character data such as the number of genes (or total number of nucleotides) should reduce stochastic error or character sampling bias (Phillips et al., 2004; Delsuc et al., 2005). The rapidly increasing amount of sequence data available to researchers from whole genomes, expressed sequence tags or cDNA libraries, and individual gene-based studies means that many analyses of these character-rich phylogenetic matrices can greatly reduce stochastic error.

This plethora of sequence data has caused some researchers to argue that large character data sets alone are sufficient to estimate an accurate phylogeny, notwithstanding the conclusions reached by numerous studies showing the importance of taxon sampling. For example, Rosenberg and Kumar (2001) conducted a simulation study indicating that adding taxa to a problematic phylogeny is less effective than adding additional characters. This paper led to a debate in the literature and reanalyses of the Rosenberg and Kumar (2001) data (Zwickl & Hillis, 2002; Pollock et al., 2002; Rosenberg & Kumar, 2003; Hillis et al., 2003). Pollock et al. (2002) reanalyzed the Rosenberg and Kumar (2001) data using a different approach to summarizing results (measurement of error), and Zwickl and Hillis (2002) re-conducted the Rosenberg and Kumar (2001) study with a different approach to study design that examined a fuller spectrum of taxon sampling strategies. Both studies concluded that taxon sampling has a very strong and positive effect on the accuracy of phylogenetic reconstruction, and showed in many cases that increasing the number of taxa had a much greater beneficial effect than increasing the number of characters. However, because of the ever-increasing availability and generation of genomic data and the difficulty of obtaining sequence data for many taxa, the debate about the relative importance of taxon sampling versus character sampling continues in the literature (Hillis et al., 2003; Rosenberg & Kumar, 2003; Rokas et al., 2003; Cummings & Meyer, 2005; Rokas et al., 2005; Hedtke et al., 2006; Gatesy et al., 2007).

More recently, Rokas et al. (2003) argued that, under parsimony, an accurate phylogeny for seven yeast taxa could be obtained using twenty genes randomly selected from a data set of 106, regardless of taxonomic sampling. The authors based this claim on the bootstrap support for each node in their tree: once twenty genes were sampled, the bootstrap support for each node rose above 95%, and the topology was identical to that of the data set using all 106 genes. Therefore, they concluded, accuracy was increased by increasing the number of genes, with the implication that many phylogenetic studies may be using an insufficient number of genes to accurately reconstruct topologies. The most obvious problem with their study design was that the authors equated high bootstrap support with accuracy (rather than more properly with repeatability). Analyzing many characters may result in convergence on a single answer, but small systematic biases can result in convergence on the incorrect answer (a point we explore in the next section).

1.2 Increased character data can reduce stochastic error, but can contribute to systematic error

Increasing the number of nucleotides will not solve inaccurate reconstruction due to method inconsistency, or systematic error. All phylogenetic methods to date have conditions in which they will perform inconsistently. Phylogenomic data sets, which tend to have large numbers of characters with relatively few, widely dispersed taxa, may be particularly prone to problems with long-branch attraction. LBA has been identified as causing spurious relationships in large-scale studies of several taxonomic groups, including mammals (Lin et al., 2002), metazoans (Anderson & Swofford, 2004; Baurain et al., 2007), arthropods (Delsuc et al., 2003) and angiosperms (Stefanovic et al., 2004). When methods are inconsistent, increasing character data can increase statistical support for an inaccurate phylogeny (Huelsenbeck & Hillis, 1993). This occurs because measures of support such as the nonparametric bootstrap proportion or Bayesian posterior probabilities are conditional on the data and the method (Delsuc et al., 2005).

A reanalysis of the Rokas et al. (2003) data set by Hedtke et al. (2006) showed that much of the conflict between genes under parsimony is due to LBA. The taxa sampled are unevenly dispersed across the tree of yeast, and a long branch leading to the outgroup has a tendency to pull particular taxa together across many independent genes. Hedtke et al. (2006) additionally used genes simulated on a more densely sampled yeast phylogeny to demonstrate that for a given

bipartition affected by long-branch attraction, bootstrap support for the wrong reconstruction increased as genes were added to the analysis (Fig. 3). However, when taxon sampling was increased, fewer genes were needed to get acceptable support for the correct bipartition. Although Hedtke et al. (2006) did not examine increases in support or accuracy across the entire tree, their results are consistent with the general finding that increased taxon sampling improves accuracy across the tree.

This discussion so far has assumed that all genes share the same evolutionary history, and that by increasing the number of nucleotides a researcher increases the signal for that one history — i.e., that topological conflicts between trees based on single genes are stochastic. However, genes do not always share an identical history, because of horizontal gene transfer, introgression, incomplete lineage sorting, or gene duplication/loss. In some cases, it may be more appropriate to analyze genes separately rather than concatenating all the data (e.g., Ane & Sanderson, 2005), particularly if a researcher is interested in identifying processes that may have affected the evolution of a species.

2 Strategies for effective taxon sampling

We have discussed how increased taxon sampling can generally improve topological estimates. Unfortunately, there is no “magic number” of taxa and genes which ensures the accuracy of a phylogeny. The percentage of taxa sampled within a taxonomic group may be more important than the total number of taxa (Yang & Goldman, 1997; Hillis, 1998). For small clades of closely-related taxa, reduced taxonomic sampling may not be problematic for phylogeny reconstruction (Poe, 1998b). Critically, whether to increase sampling within a taxonomic group of interest should not depend on whether there is statistical support for a topology, as strong support does not indicate a lack of systematic error.

2.1 Information theory and taxon addition

Several techniques have been developed from information theory to examine where taxa could be added to an analysis to increase the precision of a topological estimate (Goldman, 1998; Massingham & Goldman, 2000; Geuten et al., 2007). The observed information matrix, which is a measure of the sharpness of the curve about the likelihood function, is compared to the expected (or Fisher) information matrix when a branch is added to different parts of the

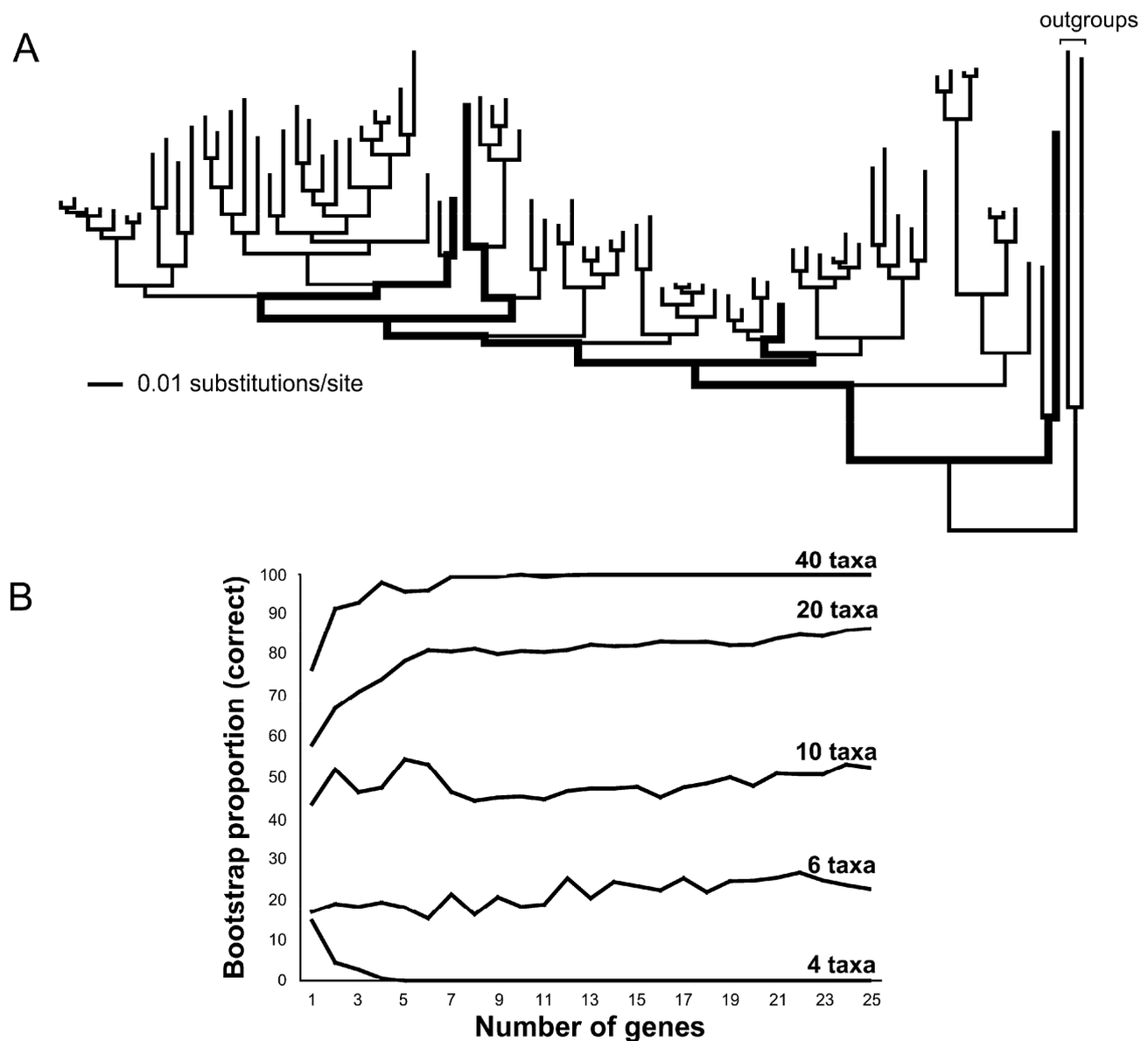


Fig. 3. **A**, Well-sampled yeast tree used for simulating data. Dark lines represent four taxa known to be susceptible to long-branch attraction under parsimony. **B**, Results of randomly sampling 1–25 genes for 4–40 taxa. As the number of genes in a parsimony analysis increases, the bootstrap for the correct reconstruction of the four-taxon statement decreases, unless taxon sampling is sufficient to break the long branches. Adapted from Hedtke et al. (2006; figs. 1 and 3).

topology (Geuten et al., 2007). In essence, this comparison indicates the increase in precision of the topology estimate that is gained by adding a taxon along a particular branch of the tree. Based on a data set from angiosperms, Geuten et al. (2007) found that information theory generally supports adding additional taxa close to where the long branch attaches to the rest of the tree, in congruence with other studies based on simulation (Graybeal, 1998; Poe & Swofford, 1999). Unfortunately, these techniques, while

promising, have not been rigorously tested, and generally assume that one has strong *a priori* expectations about where additional taxa might fall on the phylogeny.

2.2 Detecting LBA

Most studies that explore specific recommendations for increased taxon sampling focus on detecting and correcting LBA. LBA is often cited when a topology fails to meet *a priori* expectations, but caution must be applied before attributing unexpected

results to method inconsistency (Anderson & Swofford, 2004) or method bias. Many claims of LBA in published studies are *a posteriori* explanations of unexpected results. Often, when taxa are added to an analysis that originally generated an unexpected tree, topological relationships change to match expectations based on traditional taxonomy, and this change is attributed to LBA (e.g. Lin et al., 2002; Stefanovic et al., 2004; Philippe et al., 2005; Baurain et al., 2007). Here we discuss strategies which could provide useful heuristics for exploring whether method inconsistency or bias, particularly LBA, is affecting the analysis.

Disagreement among independent data sets—particularly those based on traditional morphological taxonomy versus molecular analyses—may be the first signal that LBA may be present in the data set (Lin et al., 2002; Chen et al., 2003). If two long branches are unexpectedly drawn together in one or the other analysis, LBA may be the culprit. However, a researcher must also consider the alternative that traditional taxonomy may be wrong (e.g., Ammerman & Hillis, 1992; Van Den Bussche et al., 1998). The two long branches may in fact be sister taxa, or may be brought together by LBA. To distinguish these hypotheses, one technique is to prune each long-branched taxon successively from the analysis, and observe whether the topology changes (Bergsten, 2005). Presumably, if the position of a long-branched taxon changes dependent upon the inclusion of other long-branched taxa, LBA may be implicated. Unfortunately, this test is not definitive, because other characteristics of the excluded taxa could be affecting the topology as a whole.

Simulated data generated under different hypotheses could be analyzed to compare possible topologies and get a sense of potential error rates in the analysis (Van Den Bussche et al., 1998; Sanderson et al., 2000). In this parametric bootstrapping approach, data are simulated under each hypothesis using model parameters estimated from the biological sequence data. For example, one could identify the alternative hypothesis by running a phylogenetic analysis constrained to find the best tree which does *not* place two suspect taxa as sister groups. If analyses of replicate data sets simulated on this tree tend to bring the two suspect taxa together more often than expected by chance, then one cannot reject the possibility that the two taxa are grouped due to systematic error or bias. However, as with all parametric approaches, investigators should consider the adequacy of the simulation model. Under-parameterized models may underestimate the potential for LBA.

2.3 Outgroup sampling

Outgroup taxa tend to be on long branches, either because of the processes of cladogenesis, because of extinction events between the outgroup and the ingroup, or because of inappropriate selection of outgroup taxa. When the outgroup is distantly related to the ingroup, long branches in the ingroup can be drawn toward the base of the tree by LBA, affecting ingroup relationships (Hillis, 1998; Rannala et al., 1998) or misplacing the root of the tree (Holland et al., 2003). Holland et al. (2003) used simulated data to demonstrate this effect not just for parsimony, but for maximum likelihood and distance analyses as well. Graybeal (1998) suggested that error could be reduced through the use of multiple outgroup taxa separated by short internal branches.

One method to examine whether outgroup choice is affecting the topology of the ingroup is to run the analysis using the ingroup taxa only (Holland et al., 2003; Bergsten, 2005). If the ingroup topology is influenced by the long branch leading to the outgroup, the unrooted topology with and without the outgroup will change. This technique will not allow the researcher to distinguish whether the rooting of the tree (placement of the outgroup) is correct; only whether the outgroup is influencing the ingroup topology. A second method to evaluate outgroup choice is to simulate a number of random sequences which are each used to root the tree (Sullivan & Swofford, 1997). This can indicate the relative probabilities for rooting positions when no historical signal is present in the data for the outgroup. In both cases, if use of a particular outgroup leads to LBA problems, sampling more outgroup taxa may assist in detecting homoplasy and reducing the effects of the long outgroup branch.

2.4 Ingroup sampling

If LBA is detected within the ingroup, long-branch subdivision by addition of taxa could mitigate this effect (Fig. 4; Hendy & Penny, 1989; Hillis, 1996). The strategy of long-branch subdivision has been examined using both simulated (e.g., Graybeal, 1998; Poe & Swofford, 1999; Poe, 2003) and biological (e.g., Poe, 1998a; Baurain et al., 2007) data. In parsimony analyses, there are cases in which the addition of one taxon can actually cause LBA; this has been demonstrated using simulated data in the four-taxon case when the new taxon introduces a new long branch (Rannala et al., 1998; Poe & Swofford, 1999), or when there are three long branches, and breaking one causes the remaining two to become drawn together (Fig. 4; Poe & Swofford, 1999; Poe, 2003). In this case the new taxon introduces an

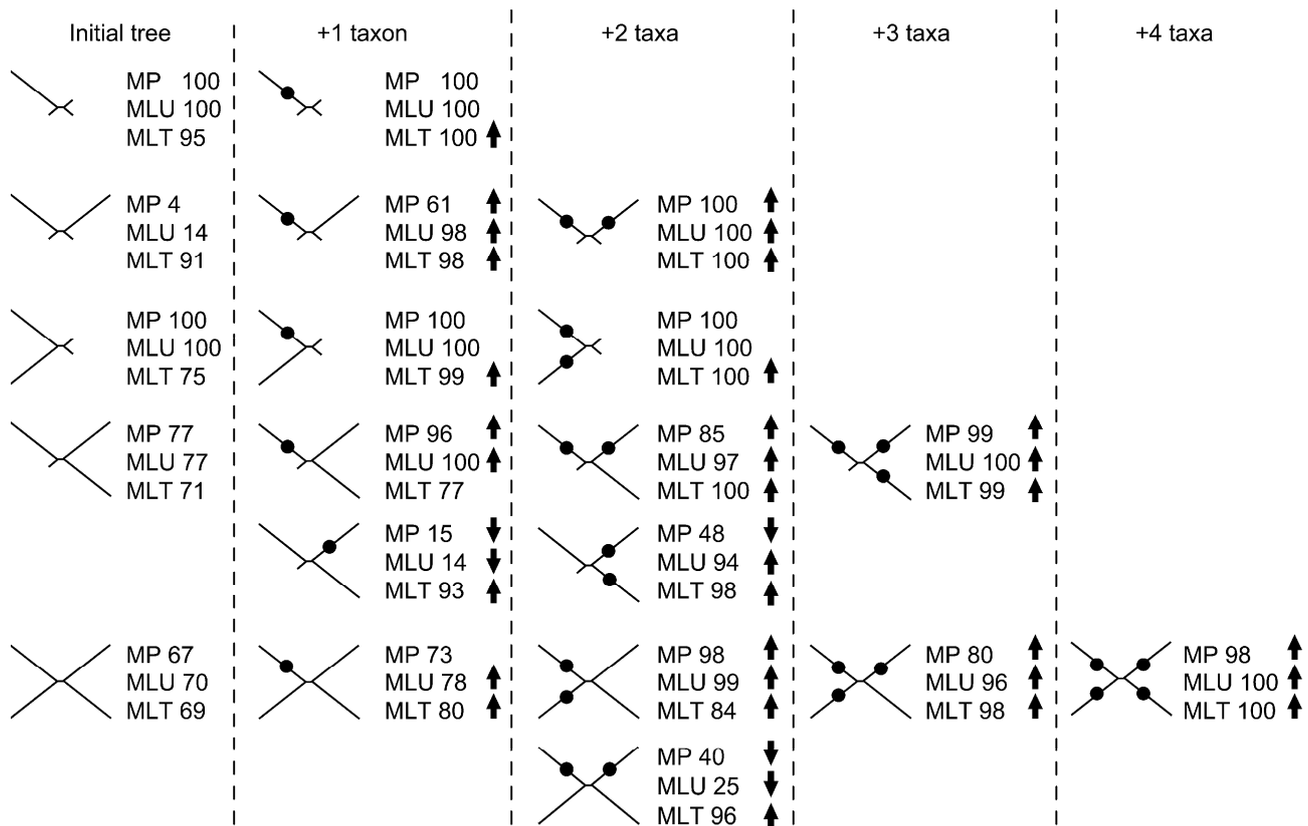


Fig. 4. Effects of long-branch subdivision on accuracy in four-taxon trees, based on simulated data. For each tree, branch lengths of long branches are 0.5, of short terminal branches 0.1, and of internal branches 0.05. Dots indicate when a long branch was broken at length 0.1 from the internal node. Poe (2003) evaluated 6 different reconstruction methods, 3 are summarized in this figure: unweighted maximum parsimony (*MP*), maximum likelihood with an under-parameterized model of substitution (*MLU*), and maximum likelihood with the true model of substitution (*MLT*). The numbers are the percentage of time the true, four-taxon tree was recovered in 100 replicate simulations. The arrows indicate increased (↑) or decreased (↓) accuracy as a result of the added taxa, and no arrow indicates that the accuracy was unaffected. Adapted from Poe (2003; fig. 3).

asymmetrical pattern of homoplasy and long-branch attraction results in an incorrect reconstruction. In addition, adding only one taxon may not be sufficient to alleviate LBA if the branch is sufficiently long (Poe, 2003; Hedtke et al., 2006) or if the added taxon is placed towards the tip (Graybeal, 1998). However, both these effects are diminished if enough taxa are added along long branches (Poe, 1998a; Anderson & Swofford, 2004; Hedtke et al., 2006). For example, in the Hedtke et al. (2006) simulation study, adding taxa increased accuracy for a particular bipartition only when the added taxa divided a long branch that was causing long-branch attraction.

2.5 Adding taxa with missing data

Adding taxa with incomplete character information to a supermatrix has primarily been evaluated in parsimony-based morphological analyses in reference to fossil data. Incomplete fossil data are not always beneficial, as they may reduce support for some nodes

(Wiens, 2003; Wiens, 2005; Cobbett et al., 2007). However, it appears that well-chosen fossil data can be helpful in breaking long branches, even if incomplete, as these morphological characters can be informative about character reconstruction at branching nodes (Donoghue et al., 1989; Huelsenbeck, 1991). The effect of missing character data on sequence analysis is still being debated, with some researchers arguing that adding taxonomic data is beneficial even if the resulting supermatrix has a large proportion of missing data (e.g., 25%: Philippe et al., 2004; 75%: Wiens & Reeder, 1995; 95%: McMahon & Sanderson, 2006). However, Lemmon et al. (in press) used simulated data to demonstrate that missing sequence data can positively mislead model-based methods. This depends in part on the relative rates of evolution for sites with and without missing data, and the topological position of those taxa with missing data. This is an area of active research, and caution should be used

when combining taxa with missing data until this issue has been more completely explored using both simulated and biological data sets.

2.6 Alternatives to increased taxon sampling

Several other techniques of combating method inconsistency have been suggested. Because inconsistency or bias results from violation of model assumptions (e.g., not accurately modeling multiple substitutions), finding a better-fitting model could solve the problem (Olsen, 1987; Whelan & Goldman, 2001; Lartillot & Philippe, 2004; Anderson & Swofford, 2004; Delsuc et al., 2005; Baurain et al., 2007; Lartillot et al., 2007). To reduce inconsistency due to LBA, it has been suggested that either eliminating fast-evolving sites from the analysis (Delsuc et al., 2002; Delsuc et al., 2005; Rodriguez-Ezpeleta et al., 2007) or coding all the data to represent only less frequent transversions ("RY" coding: Phillips et al., 2004; Delsuc et al., 2005) would reduce saturation and compositional bias in the data set, and thus reduce LBA. Using amino acid data (with models which take into account site heterogeneity, e.g., Lartillot & Philippe, 2004) may be another alternative to using raw sequence data. However, the actual effect of saturated sequences on phylogenetic analyses has been incompletely explored. Hillis (1998) simulated sequence data on a tree with such long branch lengths that the sequences would not be recognizable as homologous, but phylogenetic methods were still able to reconstruct the correct tree when taxon sampling was sufficient. Exclusion of data (whether by excluding sites or reducing their information) may be useful in eliminating the problem of a particular set of long branches, but this may be at the expense of resolution in other regions of the tree.

3 Taxon sampling affects parameter estimation

Many advances in phylogenetic analysis over the past two decades have involved model-based approaches, such as maximum likelihood and Bayesian analyses (Swofford et al., 1996; Ronquist & Huelsenbeck, 2003; Felsenstein, 2004). In general, these parametric methods outperform nonparametric methods in both simulations and experimental studies (Hillis et al., 1994a; Huelsenbeck, 1995; Cunningham et al., 1997). However, accurate phylogenetic results from model-based studies depend, at least in part, on reasonably accurate parameter estimates for the models of evolution (Goldman, 1993; Hillis et al.,

1994b; Cunningham et al., 1998; Lemmon & Moriarty, 2004; Brown & Lemmon, 2007). One of the reasons that increased taxon sampling results in more accurate phylogenetic estimation for these model-based methods is that sampling additional taxa also improves parameter estimation (Pollock et al., 1999; Sullivan et al., 1999; Pollock & Bruno, 2000; Pollock et al., 2002). In addition, as branch lengths are shortened, there are fewer unobserved changes that need to be inferred, so the accuracy of the inference becomes less dependent on the model of evolution.

In addition to their effect on phylogenetic analyses, the parameters of evolutionary models are themselves of interest to biologists. These parameters are often gene-specific, so collecting genomic-scale data from many genes across only a few taxa does little to improve our estimates of the details of evolutionary models. Instead, a thorough taxon-sampling approach is needed for each gene. Of course, the evolutionary processes may not be static across the Tree of Life for any given gene, so models that account for non-stationarity in these processes can provide better descriptions of evolutionary history (Yang & Roberts, 1995; Galtier & Gouy, 1998; Foster, 2004; Blanquart & Lartillot, 2006; Boussau & Gouy, 2006; Gowri-Shankar & Rattray, 2007; Blanquart & Lartillot, 2008; Kolaczkowski & Thornton, 2008). These models relax the assumption of time-homogeneity and can be used to detect signatures of complex evolutionary processes, such as compositional heterogeneity or heterotachy (branch-length heterogeneity), that are known to exist in biological data (Lockhart et al., 1992; Foster et al., 1997; Mooers & Holmes, 2000; Lopez et al., 2002; Jermini et al., 2004; Ane et al., 2005). Non-stationary models can greatly increase the need for even more thorough taxon sampling, because the model parameters may need to be estimated multiple times across the tree, rather than once for all taxa. It is important to note, however, that under non-stationary models, the number of parameters can increase as more sequences are added, thus increasing the computational difficulty of phylogenetic reconstruction from large data sets. Nonetheless, this obstacle may be mitigated by the use of carefully constructed priors in a Bayesian MCMC framework (Yang, 2006) and with the development of computational methods for calculating likelihoods from non-reversible models (Boussau & Gouy, 2006).

Some of the parameters that have been shown to be important for phylogenetic estimation include site-specific rates of evolutionary change; rates of change across first, second, and third positions of

codons; rates of change relative to changes in functional groups of amino-acid residues; relative rates of the various classes of transitions and transversions between nucleotide states; branch-specific rates of evolutionary change; and taxon-specific differences in base composition (Olsen, 1987; Steel et al., 1993; Hasegawa & Hashimoto, 1993; Hillis et al., 1993; Leipe et al., 1993; Goldman & Yang, 1994; Steel, 1994; Swofford et al., 1996). The number of taxa that are needed to effectively estimate these parameters differ greatly across the parameters, but all of the estimates are improved by more thorough taxon sampling. For instance, Pollock and Bruno (2000) noted significant improvement in parameter estimation (and in turn, phylogenetic estimation) as their taxon samples increased from 4 to 8 to 16 to 24 taxa. They concluded that both phylogenetic reconstruction and estimation of unknown evolutionary processes show greater improvement through increasing taxon sampling than by increasing sequence length. In some cases, reasonable parameter estimates may be obtained from external data sources, such as the HIV database, and then applied to a more limited set of taxa in the phylogenetic analysis (Hillis, 1999). However, for most taxa, the appropriate comparative data must be obtained by the investigator for a specific group of species under study.

4 Dense taxon sampling improves inferences of evolutionary processes

Beyond simply broadening our understanding of species relationships, phylogenetic trees are essential tools used in many areas of biology. Phylogenies are often used to explain broad evolutionary patterns and processes such as the evolution of adaptive traits, ancestral character states, the timing of species divergences, and variation in evolutionary rates. Many of the applications developed for these types of analyses require robust and accurate estimates of phylogeny (topology, branch lengths, and root position). This is an important consideration in and of itself; however, post-tree reconstruction applications are sensitive to reduced levels of data sampling, even when provided with an accurate phylogenetic tree.

4.1 Comparative methods

Comparative analyses are a fundamental component in the fields of evolutionary biology, behavior, and ecology. The development of statistical methods that incorporate phylogenetic trees (Felsenstein, 1985) have allowed for robust and reliable tests of the

evolution of adaptive traits and the processes that might drive diversification. For example, these methods have been used to reveal patterns in the biodiversity of marine teleost fishes (Alfaro et al., 2007) and to show that independent origins of dietary specialization have been a major factor in the evolution of defensive mechanisms in neotropical poison frogs (Darst et al., 2005). Comparative analyses of character evolution using phylogenetic comparative methods require attention to adequate sampling at many levels. At the intraspecific level, poor sampling of organismal attributes can lead to measurement error, which may result in an underestimation of the variance of contrasts between sister taxa (Ricklefs & Starck, 1996). Generation of a robust phylogeny is extremely important since different comparative methods have different ways of dealing with topological uncertainty (Purvis et al., 1994). In addition, fewer taxa (and thus fewer internal nodes for calculating contrasts) can lead to increased variance and uncertainty in the results. Ackerly (2000) used simulated data to show that the statistical power of several comparative tests decreased as the sample size of taxa decreased, and that careful attention should be paid to how species are sampled for these analyses. Biased taxon sampling, particularly with respect to the characters of interest, can lead to systematic biases in the calculation of statistical correlations between characters. The results presented by Ackerly (2000) indicate that uniform, random sampling of taxa does not introduce error in phylogenetic comparative methods.

4.2 Ancestral character states

An integral component of phylogenetic comparative analyses and other evolutionary applications is the reconstruction of ancestral character states. These methods use phylogenetic trees and branch lengths to infer the states of discrete or continuous characters at ancestral nodes, and have been used to reconstruct such diverse ancestral characters as the advertisement calls of frogs in the genus *Physalaemus* (Ryan & Rand, 1995, 1998), the fruiting-body forms of homobasidiomycetes (Hibbett, 2004), and ancient bacterial protein sequences (Gaucher et al., 2003). Dense taxon sampling is also an important consideration for ancestral-state reconstruction methods. Salisbury and Kim's (2001) analyses of simulated data and trees indicated that the accuracy of parsimony ancestral-state estimation decreases with reduced taxon sampling and increased rates of character evolution (Fig. 5). Because parsimony methods do not account for unobserved changes, they usually underestimate the number of changes along a branch (Fitch & Bruschi, 1987;

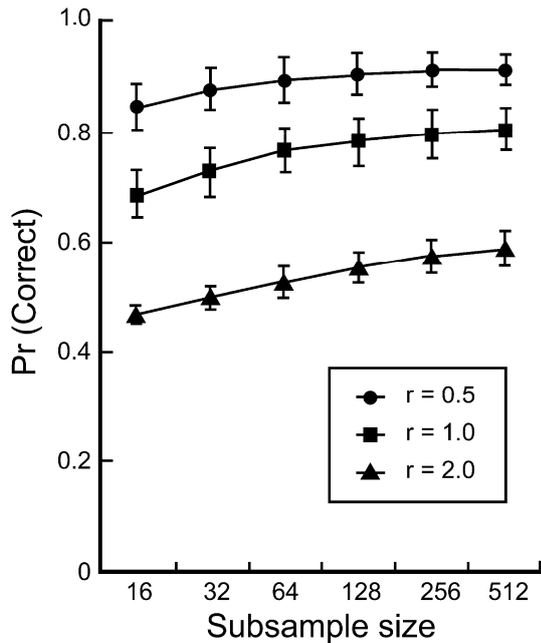


Fig. 5. The mean probabilities, $Pr(\text{Correct})$, of correctly estimating the root state of a binary character evolving at 3 different rates (r) on subsamples of 512-taxon, pure-birth model tree topologies. Each point is the mean for a sample of 100 trees and the error bars represent the ± 1 standard deviation. Adapted from Salisbury and Kim (2001; fig. 1).

Fitch & Beintema, 1990; Huelsenbeck & Lander, 2003). Dense taxon sampling can reduce this effect and improve the accuracy of parsimony ancestral-state estimates. Maximum likelihood and Bayesian methods for reconstructing ancestral states have also been developed (Pagel, 1994; Schluter et al., 1997; Pagel, 1999; Huelsenbeck & Bollback, 2001; Pagel et al., 2004). These parametric ancestral-state reconstruction methods are also sensitive to high rates of character evolution. However, Schluter et al. (1997) showed that parsimony ancestral-state reconstruction methods often fail to identify ambiguous-node state estimates. Conversely, maximum likelihood and Bayesian methods are less likely to provide misleading results because these methods incorporate branch-length information and explicit models of character evolution and quantify uncertainty in ancestral-state estimates (provided that the model assumptions are adequate). Bayesian approaches, in particular, use Markov chain Monte Carlo sampling to accommodate and quantify uncertainty in the tree topology, branch lengths, ancestral states, and model parameters (Huelsenbeck & Bollback, 2001; Pagel et al., 2004). Denser taxon sampling reduces the number of unobserved evolu-

tionary events, and so is also expected to simplify and improve the reconstruction of ancestral states in model-based analyses.

4.3 Divergence time estimation

A primary field of research in evolutionary biology involves estimation of the timing and rate of evolutionary processes. In these applications, phylogenetic trees are used to date speciation events and infer lineage-specific substitution rates. Reliable estimates of species divergence times are fundamental components for understanding historical biogeography, testing hypotheses of adaptive character evolution, and estimating speciation and extinction rates. However, divergence time estimation is hindered by the fact that the rate of evolution and time are intrinsically linked when inferring genetic distances between lineages. Several methods test for variation in the rates of molecular evolution or tease apart the rate of substitution and time by applying models for estimating lineage-specific substitution rates. These methods include strict molecular clock models (Zuckerkandl & Pauling, 1962; Langley & Fitch, 1974), local molecular clocks (Kishino & Hasegawa, 1990; Rambaut & Bromham, 1998; Yoder & Yang, 2000; Yang & Yoder, 2003), non-parametric and semi-parametric methods for estimating autocorrelated substitution rates (Sanderson, 1997, 2002), and Bayesian methods for estimating autocorrelated and uncorrelated rates (Thorne et al., 1998; Huelsenbeck et al., 2000; Kishino et al., 2001; Thorne & Kishino, 2002; Drummond et al., 2006; Lepage et al., 2006). These various approaches have been applied to a number of biological data sets (e.g., Yang & Yoder, 2003; Smith et al., 2006; Bell, 2007; Hugall et al., 2007; Roelants et al., 2007; Zhou & Holmes, 2007). Current implementations of most of these methods require a fixed tree topology and sometimes fixed branch lengths (Thorne & Kishino, 2002; Sanderson, 2003; Lepage et al., 2007; for exceptions see Drummond et al., 2006). Because of their reliance on phylogenetic data, these methods can be sensitive to taxon sampling density. Robinson et al. (1998) evaluated the effect of reduced taxon sampling on the performance of the relative-rates test. The relative-rates test (Sarich & Wilson, 1973; Wu & Li, 1985) is used to compare the substitution rates between two species and has been extended for analyzing larger phylogenetic trees to detect rate variation (Li & Bousquet, 1992; Takezaki et al., 1995). The simulation study of Robinson et al. (1998) showed that increased proportions of taxon sampling improved the accuracy of the relative-rates test.

Most of the work exploring the accuracy of molecular dating methods has revealed that these methods are very sensitive to the fossil calibrations used and little is known about the impact of taxon sampling on divergence time estimates (Yang & Rannala, 2006; Rutschmann et al., 2007; Hugall et al., 2007). A recent study by Hug and Roger (2007) used two biological data sets with low levels of taxon sampling (30 metazoan taxa with two outgroup species, and a 36 taxon data set that spanned all eukaryotes) and concluded that, for these data sets, reduced taxon sampling was not an important factor in the estimation of node times. However, their analyses showed that the choice and application of fossil calibration points resulted in a significant impact on the estimates of node ages. From their results, Hug and Roger (2007) recommended that biologists should focus on improving the number and quality of their fossil calibrations and not on increasing taxon sampling, provided there are enough taxa to obtain a reliable estimate of phylogeny. However, because of the sparsely sampled data sets used in this study and the demonstrated extreme sensitivity of these data to fossil constraints, Hug and Roger's (2007) results may not apply to a more general set of conditions and the importance of dense taxon sampling for estimating species divergence times is still an open question.

Node-density effects, as a result of uneven taxon sampling, may adversely affect molecular dating analyses (Hugall & Lee, 2007). Based on the studies demonstrating the sensitivity of divergence time estimation methods to fossil calibration choice (Near & Sanderson, 2004; Near et al., 2005; Roger & Hug, 2006; Yang & Rannala, 2006; Ho, 2007; Hugall et al., 2007; Rutschmann et al., 2007), together with studies emphasizing the importance of increased taxon sampling on phylogenetic reconstruction methods and the estimation of evolutionary parameters (Lecointre et al., 1993; Hillis, 1996, 1998; Graybeal, 1998; Rannala et al., 1998; Pollock & Bruno, 2000; Zwickl & Hillis, 2002; Pollock et al., 2002; Poe, 2003; DeBry, 2005; Hedtke et al., 2006), we recommend increased collection of fossils and improved taxon sampling density for these types of analyses, whenever possible. Maximizing the number of fossil calibration points goes hand-in-hand with increasing taxon sampling because densely sampled trees provide a greater number of internal nodes on which an investigator can place a fossil calibration. Moreover, investigators are far more restricted by the availability of fossils and other types of information for calibrating divergences than by the availability of extant taxa. Further investigation using

simulations and well-sampled data sets of living and fossil taxa should help shed light on this issue. Because extensive taxon sampling (especially of fossil taxa) is sometimes impractical, Bayesian methods for divergence time estimation present promising opportunities to account for uncertainty in phylogenies by simultaneously estimating the tree topology and branching times (Drummond et al., 2006). These methods can also incorporate information on taxon sampling density in the form of priors on the distribution of divergence times (Yang & Rannala, 1997, 2006).

4.4 Evaluating diversification rates

Phylogenetic trees are fundamental for understanding variation in species diversity. Methods for elucidating patterns of speciation and extinction measure the shape of phylogenies to detect shifts in diversification rates or to estimate global net diversification rates. Phylogenetic tree shape can be measured by quantifying how node ages are distributed over time or by calculating the degree of asymmetry among lineages in the tree. Measures of tree shape can be compared to a null model that assumes all lineages have experienced the same rate of diversification (Shao & Sokal, 1990; Kirkpatrick & Slatkin, 1993; Nee et al., 1994b; Pybus & Harvey, 2000; Agapow & Purvis, 2002). Analyses of the temporal distribution of diversification events use branch lengths obtained from time-adjusted phylogenies to estimate and detect large shifts in speciation and extinction rates (Nee et al., 1994b; Pybus & Harvey, 2000). For example, Becerra (2005) applied these methods to investigate temporal and biogeographic processes that may have shaped the diversity of the plant genus *Bursera*. The results of this study indicate that the radiation of this group is associated with the establishment of tropical dry forest habitat in Mexico. However, inadequate taxon sampling has a significant impact on these methods (Nee et al., 1994a). Nee et al. (1994a) used lineages-by-time plots to show that incomplete taxon sampling can result in an apparent reduction in the rate of diversification over time, even when the tree evolved under constant rates of speciation and extinction.

Analyses based on topology measure asymmetry in the distribution of lineages over a tree to test for changes in diversification rates. These methods evaluate the balance either at a single node or over the entire tree (Shao & Sokal, 1990; Kirkpatrick & Slatkin, 1993; Agapow & Purvis, 2002) and are often used to detect patterns characteristic of rapid radiations in phylogenetic trees (Guyer & Slowinski, 1993; Chan &

Moore, 1999, 2002). The degree of taxon sampling is an important consideration when conducting these analyses. Several studies have shown that published phylogenies are (on average) much more imbalanced than expected under a model assuming constant diversification rates (Guyer & Slowinski, 1991; Heard, 1992; Mooers, 1995; Purvis & Agapow, 2002; Holman, 2005; Blum & François, 2006; Heath et al., 2008). Mooers (1995) compared the level of tree imbalance in a collection of published phylogenies and found that incomplete trees are more imbalanced than completely sampled phylogenies. Another study by Heath et al. (2008) examined the effect of random taxon sampling on empirical trees and on phylogenies simulated under different models of cladogenesis. They found that reduced taxon sampling of empirical trees and trees simulated under variable and autocorrelated speciation and extinction rates causes an increase in node imbalance. These results suggest that poor taxon sampling leads to an increase in the apparent rate variation because of the overrepresentation of older nodes. The bias caused by incomplete species sampling must be considered when using phylogenies to test hypotheses about species diversity.

5 Innovations in reconstruction algorithms and the analysis of large data sets

Until recently, computational constraints on phylogenetic analyses made inclusion of large numbers of taxa impractical for many biologists. However, developments in computational power, parallel computation, and phylogenetic algorithms have greatly decreased computational constraints for phylogenetic analyses of many taxa, even for the most computationally demanding parametric approaches (e.g., Brauer et al., 2002; Lemmon & Milinkovitch, 2002; Guindon & Gascuel, 2003; Ronquist & Huelsenbeck, 2003; Stamatakis, 2006; Minh et al., 2005; Zwickl, 2006). Quick but imprecise clustering techniques, such as the widely used neighbor-joining algorithm (Saitou & Nei, 1987), are rapidly being replaced by methods that more thoroughly explore solution space using a clearly defined model of evolution (implemented in such programs as RAXML, GARLI, PhyML, IQPNNI, MRBAYES, and PAUP*; see <http://evolution.genetics.washington.edu/phylip/software.html> for more information). Analyses of hundreds to thousands of taxa have become routine for parsimony, maximum likelihood, and Bayesian approaches, and analyses of tens-of-thousands of taxa are now feasible, even for parametric methods. Therefore, computa-

tional constraints can no longer be viewed as a serious impediment to thorough taxon sampling. Instead, the limitations of taxon sampling have now shifted to problems of taxon availability and the constraints of specimen and data collection. Given the many benefits of thorough taxon sampling summarized in this paper, we advise biologists to carefully consider taxon sampling design in planning, conducting, and interpreting phylogenetic analyses. In many cases, increasing taxon sampling is one of the most practical and beneficial approaches to increasing the accuracy of phylogenetic estimates and the biological inferences that are derived from phylogenetic trees.

Acknowledgements We thank professors Deyuan Hong, Zhiduan Chen, Chengxin Fu, Michael J. Donoghue, and Yin-Long Qiu for hosting the symposium on “Evolutionary Biology in the 21st Century—Tracing Patterns of Evolution through the Tree of Life,” and for the invitation to D.M.H. to present this paper. The symposium was supported by a grant from the National Natural Science Foundation of China. A.J. Abrams, J. Brown, M. Morgan, G. Pauly, R. Springman, M. Swenson, R. Symula, D. Zwickl, Hervé Philippe, and an anonymous reviewer provided helpful comments on this manuscript. D.M.H. gratefully acknowledges grant support from the United States National Science Foundation (NSF). T.A.H. was funded by a graduate research traineeship provided by an NSF IGERT grant in Computational Phylogenetics and Applications to Biology awarded to the University of Texas, Austin. S.M.H. was supported by an NSF pre-doctoral fellowship.

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